COMMUNICATION

An efficient protocol for incorporation of an unnatural amino acid in perdeuterated recombinant proteins using glucose-based media

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Abstract The in vivo incorporation of unnatural amino acids into proteins is a well-established technique requiring an orthogonal tRNA/aminoacyl-tRNA synthetase pair specific for the unnatural amino acid that is incorporated at a position encoded by a TAG amber codon. Although this technology provides unique opportunities to engineer protein structures, poor protein yields are usually obtained in deuterated media, hampering its application in the protein NMR field. Here, we describe a novel protocol for incorporating unnatural amino acids into fully deuterated proteins using glucose-based media (which are relevant to the production, for example, of amino acid-specific methyllabeled proteins used in the study of large molecular weight systems). The method consists of pre-induction of the pEVOL plasmid encoding the tRNA/aminoacyl-tRNA synthetase pair in a rich, H₂O-based medium prior to exchanging the culture into a D₂O-based medium. Our protocol results in high level of isotopic incorporation $(\sim 95\%)$ and retains the high expression level of the target protein observed in Luria-Bertani medium.

Keywords Site-specific labeling · Isotopic labeling · Unnatural amino acids · Spin-labeling

In vivo incorporation of unnatural amino acids into proteins (Wang et al. 2001; Liu et al. 2007) has significantly expanded frontiers in protein engineering. To date, over 50 unnatural amino acids have been genetically encoded (Liu

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and Schultz 2006; Xie and Schultz 2006; Brustad et al. 2008a, b; Cellitti et al. 2008), permitting the manipulation of physicochemical, biological, and pharmacological properties of proteins with exquisite control over structure.

The advantages of creating proteins with unnatural amino acids for NMR studies have been recently discussed (Jones et al. 2010). For example, the introduction of unnatural amino acids bearing post-translational modifications (Liu and Schultz 2006; Xie et al. 2007; Guo et al. 2008) can be particularly important for structural and dynamic investigations of signaling pathways. In addition, site-specific incorporation of NMR active (¹H, ¹³C, ¹⁵N and ¹⁹F) unnatural amino acids into silent backgrounds dramatically reduces spectral complexity, thereby facilitating the study of large proteins by in-vitro and in-vivo NMR techniques (Cellitti et al. 2008; Jones et al. 2010; Lampe et al. 2010; Li et al. 2010; Watt et al. 2011). Particularly attractive are methods that employ unnatural amino acids to design binding sites for paramagnetic metal ions (Lee and Schultz 2008; Lee et al. 2009) or nitroxide spin-labels (Fleissner et al. 2009). Indeed, incorporation of unnatural amino acids having bio-orthogonal chemical functionality in the context of the native protein can facilitate production of site-specific, paramagnetically-labeled samples, critical for the characterization of transient, sparsely-populated states (Clore and Iwahara 2009) and very valuable for resonance assignment (Pintacuda et al. 2004; Venditti et al. 2011) in large proteins and protein complexes.

Incorporation of unnatural amino acids is accomplished by co-expressing a bio-orthogonal tRNA/aminoacyl-tRNA synthetase pair specific for the relevant unnatural amino acid together with the protein of interest whose DNA sequence has been modified by substitution of a TAG amber codon at the site of desired incorporation (Wang et al. 2001). However, lower protein yields are generally



obtained by this method owing to translation termination. Thus, to reduce truncation problems, an optimized E. coli incorporation system consisting of a plasmid that overexpresses the aminoacyl-tRNA synthetase at the time of induction has been developed (Cellitti et al. 2008), and the effects of the degree, timing, and temperature of induction have been extensively tested (Young et al. 2010). These studies demonstrated that the simultaneous overexpression of the aminoacyl-tRNA synthetase and the target protein at 30°C dramatically reduces truncation problems, enabling production of sufficient quantities of protein for NMR studies using minimal or rich media (Cellitti et al. 2008; Young et al. 2010). More recently, the expression of the protein FAS-TE incorporating the unnatural amino acid p-methoxy-phenylalanine (p-OMePhe) in deuterated M9 medium supplemented with 0.5% d-glycerol and 0.1 mM FeCl₃ has been also reported (Jones et al. 2010). However, the yield of the protein was modest ($\sim 25\%$ of the level obtained in rich medium) and, to the best of our knowledge, no other successful incorporation of unnatural amino acids into deuterated proteins has been reported to date.

Here, we investigate the expression levels for a mutant (C575A) of the C-terminal domain (A261-C575) of the E. coli enzyme I (EIC) of the bacterial phosphotransfer system (PTS), incorporating the unnatural amino acid p-acetyl-L-phenylalanine (pAcF) at position M469 in protonated and deuterated media (Table 1). pAcF is particularly relevant to the NMR community since it reacts quantitatively with a new hydroxylamine functionalized nitroxide spin-label, known as HO-4120 (Fleissner et al. 2009), enabling production of site-directed, paramagnetically-labeled proteins even in the presence of surface exposed Cys residues which interfere with standard sitedirected spin-labeling techniques. We focused our attention on glucose-based deuterated media since these are routinely employed for the production of amino acid-specific, methyl labeled proteins (Tugarinov et al. 2006) that are extremely useful in NMR studies of high molecular weight systems (Mittermaier and Kay 2006).

We initially followed the optimized protocol for incorporation of pAcF into proteins using a protonated medium (Young et al. 2010). Briefly, BL21 star (DE3) E. coli cells were co-transformed with pET11a/EIC (containing the desired TAG mutant) and the pEVOL plasmid that contains the orthogonal tRNA and aminoacyl-tRNA synthetase specific for pAcF (Young et al. 2010). Transformed bacteria were then plated onto an LB-agar plate containing ampicillin (100 µg/ml) and chloramphenicol (34 µg/ml) for selection. A single colony was used to inoculate 50 ml of Luria-Bertani (LB) medium containing the aforementioned antibiotics and grown overnight at 37°C. The following day the starter culture was diluted to $A_{600} \sim 0.05$ in 50 ml of each medium listed in Table 1 and grown at 37°C to $A_{600} \sim 0.6$. pAcF was then added to each culture to a final concentration of 2 mM. After 1 h, the cultures were moved to 30°C and the pET and pEVOL plasmids were induced with 1 mM isopropyl-β-D-1-thiogalactopyranoside (IPTG) and 0.02% L-arabinose, respectively. After 12 h, 1 ml of each culture was harvested by centrifugation, resuspended in 300 µl of B-PER (Pierce) and the soluble fraction was checked by SDS-page.

Thick overexpression bands are clearly observed for both the full length and truncated proteins in LB medium (Fig. 1A). In contrast, only a small amount of truncated

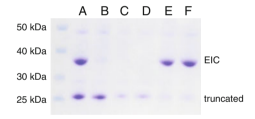


Fig. 1 Expression of EIC M469pAcF/C575A in different media. *Lanes A, B, C* and *D* correspond to expression in LB, M9^{H2O}, M9^{D2O} and M9^{ISO} media (see Table 1), respectively. *Lanes E* and *F* correspond to expression performed by pre-inducing the pEVOL plasmid in LB medium prior to moving the culture to the deuterated M9^{D2O} and M9^{ISO} media, respectively. The left-hand most lane displays standard molecular weight markers

Table 1 Composition of the various culture media used in the present work

Medium	Composition (per 1 l)
LB	10.0 g bacto-tryptone, 5.0 g yeast extract, 5.0 g NaCl, 200 μl 5 M NaOH
M9 ^{H2O}	6.5 g Na ₂ HPO ₄ , 3.0 g KH ₂ PO ₄ , 0.5 g NaCl, 1.0 g ¹⁵ NH ₄ Cl, 2.0 g D-glucose, 120 mg MgSO ₄ , 11 mg CaCl ₂ , 10 ml 100X MEM vitamins mix (VWR)
M9 ^{D2O}	$6.5~g~Na_2HPO_4,~3.0~g~KH_2PO_4,~0.5~g~NaCl,~1.0~g~^{15}NH_4Cl,~2.0~g~_D-glucose-1,2,3,4,5,6,6-d_7,~120~mg~MgSO_4,~11~mg~CaCl_2,~10~ml~100X~MEM~vitamins~mix~(VWR)$
M9 ^{ISO}	$6.5~g~Na_2HPO_4,~3.0~g~KH_2PO_4,~0.5~g~NaCl,~1.0~g~^{15}NH_4Cl,~2.0~g~D-glucose-1,2,3,4,5,6,6-d_7,~120~mg~MgSO_4,~11~mg~CaCl_2,~10~ml~100X~MEM~vitamins~mix~(VWR),~1.0~g~^{15}N-^2H~IsoGro~(Isotec)$

The LB and $M9^{\rm H2O}$ media were prepared in water, while $M9^{\rm D2O}$ and $M9^{\rm ISO}$ media were prepared in 99.9% D_2O . The MEM vitamin mix used in the deuterated media was lyophilized and resuspended in 99.99% D_2O before usage

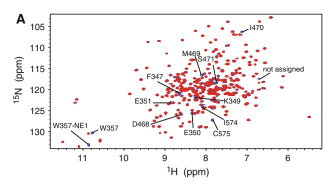


Table 2 Final optical density (A_{600}) and yield of EIC M469pAcF/C575A measured for the 50 ml test cultures

Medium	Final A ₆₀₀	Yield (mg) ^a
LB	~2.4	~9.1
M9 ^{H2O}	~ 2.0	_
$M9^{D2O}$	~1.5	_
M9 ^{ISO}	~1.6	_
LB-M9 ^{D2O}	~ 2.0	~8.4
LB-M9 ^{ISO}	~2.2	~10.2

LB-M9^{D2O} and LB-M9^{ISO} indicate cultures pre-induced in LB medium with L-arabinose and then moved to deuterated media (see text) ^a Measured for purified EIC M469pAcF/C575A (residues 261–575) using an extinction coefficient $\varepsilon_{280} = 19.94 \text{ mM}^{-1} \text{ cm}^{-1}$. In all cases conversion to mg units was carried out by using the theoretical mass for the EIC domain at natural abundance (35,206 Da)

EIC with no detectable band for the full-length protein was obtained from cultures in protonated and deuterated minimal media (Fig. 1B-D). This finding cannot be ascribed to an intrinsically low expression of EIC in minimal media since ~ 15 mg of U-[2 H- 15 N] wild type EIC was obtained from 50 ml cultures in M9^{D2O} (Table 1). An alternative explanation may be the reduced expression of the tRNA/ aminoacyl-tRNA synthetase pair which is under control of the ara promoter and could be largely inhibited at the high glucose concentration used in the tested minimal media. To test this hypothesis the starter culture was diluted to $A_{600} \sim 0.05$ in 50 ml of LB medium and grown at 37°C to an $A_{600} \sim 0.4$. The pEVOL plasmid was then induced with 0.02% L-arabinose for 2 h, while still in protonated, LB medium. Bacteria were collected by gentle centrifugation (170 g for 25 min) and resuspended in 50 ml of two



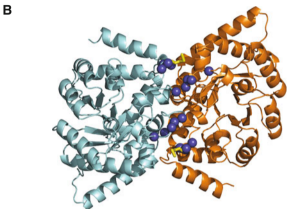


Fig. 2 A 600 MHz ¹H-¹⁵N TROSY spectra of wild-type (*blue*) and M469*p*AcF/C575A (*red*) EIC. The NMR spectra were acquired at 37°C. Sample conditions were as follow: 400 μM EIC, 20 mM Tris, pH 7.4, 100 mM NaCl, 4 mM MgCl₂, 1 mM EDTA. Under these conditions EIC is present as a 70 kDa dimer (Patel et al. 2006). Resonance assignment for cross-peaks showing large chemical shift differences in the two EIC constructs is reported on the wild-type spectrum. **B** X-ray structure of the wild-type EIC dimer (PDB code 2HWG) (Teplyakov et al. 2006). The monomeric subunits are colored in cyan and orange. Residues whose resonance show large combined ¹H_N/¹⁵N chemical shift differences (>0.1 ppm) as a consequence of the M469*p*AcF and C575A mutations are indicated as *blue spheres*. M469 is shown as *yellow sticks*

Table 3 LC-MS analysis of the natural abundance and U-[2H,15N]-labeled EIC M469pAcF/C575A samples

Medium	Theoretical mass (Da) ^a	Observed mass (Da)	Theoretical number of isotopic substitutions ^b	Observed number of isotopic substitutions ^c	Isotopic incorporation (%)
LB	35,206	35,205	-	_	_
LB-M9 ^{D2O}	37,551	36,428	2,345	2,223	94.8
LB-M9 ^{ISO}	37,551	37,441	2,345	2,236	95.4
WT-M9 ^{D2O}	37,547	37,530	2,368 ^d	2,351 ^d	99.3

Data were acquired under denaturing conditions with a liquid chromatography (Zorbax 300SB-C3 column) mass spectrometry electrospray ionization time-of-flight (LC-MS ESI-TOF) instrument (HP 1100 HPLC/MSD). Protein samples (6 μ M final concentration) were diluted in water and acidified with 10% acetic acid. Mass spectra were recorded in the positive ion mode (mass range 500–2,000) and results were analyzed using the HP ChemStation software. In the last row (WT-M9^{D2O}) LC-MS data for U-[2 H, 15 N]-labeled wild-type (WT) EIC expressed in the M9^{D2O} medium are reported



^a Theoretical masses were calculated assuming 100% isotopic incorporation and complete back exchange to ¹H at labile hydrogen positions

^b This is the expected number of ¹⁵N and ²H atoms incorporated into EIC and is calculated by taking the difference between the theoretical mass in LB and deuterated media

^c This is the observed number of ¹⁵N and ²H atoms incorporated into EIC and is calculated by taking the difference between the observed mass in LB and deuterated media

^d Calculated using the theoretical mass in LB for the natural abundance wild-type EIC (35,179 Da)

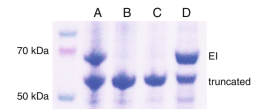


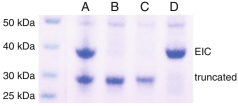
Fig. 3 Expression of full-length (residues 1-575) EI M469 pAcF/C575A (*left panel*) and EIC A494pAcF/C575A (*right panel*) in different media. *Lanes A*, B and C correspond to expression in LB, M9^{H2O} and M9^{ISO} media (see Table 1), respectively. *Lane D*

different recipes of deuterated media, $M9^{D2O}$ and $M9^{ISO}$ (Table 1), to a final A_{600} of ~ 0.8 . Subsequently, 0.5 g/l pAcF (final concentration 2 mM) and 0.02% L-arabinose were added to each culture. After 1 h of incubation at 30°C the expression of EIC was induced with 1 mM IPTG and the two cultures were shaken at 30°C for 12 h. In stark contrast to the near zero yield afforded by the standard protocol, a large amount of full-length protein was obtained using the latter protocol, with yields comparable to the one obtained in LB medium (Fig. 1E, F; Table 2). These data clearly indicate that pre-inducing the pEVOL plasmid in LB medium is extremely beneficial for protein expression.

The degree of isotopic incorporation in the proposed protocol was tested by liquid chromatography/mass spectrometry on purified EIC M469pAcF/C575A. The results are summarized in Table 3, demonstrating that high levels of isotopic incorporation (\sim 95%) are obtained in both the deuterated media tested.

As a final quality test on the produced protein, a comparison of the ¹H-¹⁵N TROSY spectra of the wild type and mutant proteins is reported in Fig. 2. The two spectra show excellent overlap (Fig. 2A), differing only in cross-peaks assigned to residues in the vicinity of the introduced mutation (Fig. 2B), indicating that the introduction of the unnatural amino acid *p*AcF at position 469 does not alter the overall folding of EIC.

In conclusion, we have presented a new protocol for incorporating the unnatural amino acid *p*AcF into deuterated EIC using the recently developed pEVOL vector (Young et al. 2010). We have shown that pre-inducing the expression of the tRNA/aminoacyl-tRNA synthetase pair in LB medium prior to switching the culture to a deuterated, glucose-based medium results in protein yields comparable to that obtained in rich medium, with no significant effects on the isotopic incorporation level. Similar results were also obtained for an EIC construct incorporating *p*AcF at position A494, as well as for the full-length EI (subunit molecular weight of 64 kDa) incorporating *p*AcF at position M469 (Fig. 3), suggesting that our procedure is of general applicability. We therefore fully expect the overall protocol presented here to be applicable to other unnatural



corresponds to expression performed by pre-inducing the pEVOL plasmid in LB medium prior to moving the culture to the deuterated M9^{ISO} medium. The left-hand most lane in each panel displays standard molecular weight markers

amino acids and protein constructs. It is worth noting, however, that the reduced expression levels obtained with standard protocols is probably related to the high glucose content of the tested minimal media. Thus, pre-induction of the pEVOL plasmid in LB media may not be needed when using other carbon sources.

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References

Brustad E, Bushey ML, Brock A, Chittuluru J, Schultz PG (2008a) A promiscuous aminoacyl-tRNA synthetase that incorporates cysteine, methionine, and alanine homologs into proteins. Bioorg Med Chem Lett 18:6004–6006

Brustad E, Bushey ML, Lee JW, Groff D, Liu W et al (2008b) A genetically encoded boronate-containing amino acid. Angew Chem Int Ed Engl 47:8220–8823

Cellitti SE, Jones DH, Lagpacan L, Hao X, Zhang Q et al (2008) In vivo incorporation of unnatural amino acids to probe structure, dynamics, and ligand binding in a large protein by nuclear magnetic resonance spectroscopy. J Am Chem Soc 130:9268–9281

Clore GM, Iwahara J (2009) Theory, practice, and applications of paramagnetic relaxation enhancement for the characterization of transient low-population states of biological macromolecules and their complexes. Chem Rev 109:4108–4139

Fleissner MR, Brustad EM, Kalai T, Altenbach C, Cascio D et al (2009) Site-directed spin labeling of a genetically encoded unnatural amino acid. Proc Natl Acad Sci USA 106:21637–21642

Guo J, Wang J, Lee JS, Schultz PG (2008) Site-specific incorporation of methyl- and acetyl-lysine analogues into recombinant proteins. Angew Chem Int Ed Engl 47:6399–6401

Jones DH, Cellitti SE, Hao X, Zhang Q, Jahnz M et al (2010) Sitespecific labeling of proteins with NMR-active unnatural amino acids. J Biomol NMR 46:89–100

Lampe JN, Brandman R, Sivaramakrishnan S, de Montellano PR (2010) Two-dimensional NMR and all-atom molecular dynamics of cytochrome P450 CYP119 reveal hidden conformational substates. J Biol Chem 285:9594–9603

Lee HS, Schultz PG (2008) Biosynthesis of a site-specific DNA cleaving protein. J Am Chem Soc 130:13194–13195



- Lee HS, Spraggon G, Schultz PG, Wang F (2009) Genetic incorporation of a metal-ion chelating amino acid into proteins as a biophysical probe. J Am Chem Soc 131:2481–2483
- Li C, Wang GF, Wang Y, Creager-Allen R, Lutz EA et al (2010) Protein (19)F NMR in Escherichia coli. J Am Chem Soc 132:321–327
- Liu CC, Schultz PG (2006) Recombinant expression of selectively sulfated proteins in *Escherichia coli*. Nat Biotechnol 24: 1436–1440
- Liu W, Brock A, Chen S, Chen S, Schultz PG (2007) Genetic incorporation of unnatural amino acids into proteins in mammalian cells. Nat Methods 4:239–244
- Mittermaier A, Kay LE (2006) New tools provide new insights in NMR studies of protein dynamics. Science 312:224–228
- Patel HV, Vyas KA, Mattoo RL, Southworth M, Perler FB et al (2006) Properties of the C-terminal domain of Enzyme I of the Escherichia Coli phosphotransferase system. J Biol Chem 281:17579–17587
- Pintacuda G, Keniry MA, Huber T, Park AY, Dixon NE et al (2004) Fast structure-based assignment of ¹⁵N HSQC spectra of selectively ¹⁵N-labeled paramagnetic proteins. J Am Chem Soc 126:2963–2970
- Teplyakov A, Lim K, Zhu PP, Kapadia G, Chen CC et al (2006) Structure of phosphorylated enzyme I, the phosphoenolpyruvate:sugar phosphotransferase system sugar translocation signal protein. Proc Natl Acad Sci USA 103:16218–16223

- Tugarinov V, Kanelis V, Kay LE (2006) Isotope labeling strategies for the study of high-molecular-weight proteins by solution NMR spectroscopy. Nat Protoc 1:749–754
- Venditti V, Fawzi NL, Clore GM (2011) Automated sequence- and stereo-specific assignment of methyl-labeled proteins by paramagnetic relaxation and methyl-methyl nuclear overhauser enhancement spectroscopy. J Biomol NMR 51:319–328
- Wang L, Brock A, Herberich B, Schultz PG (2001) Expanding the genetic code of *Escherichia coli*. Science 292:498–500
- Watt ED, Rivalta I, Whittier SK, Batista VS, Loria JP (2011) Reengineering rate-limiting, millisecond enzyme motions by introduction of an unnatural amino acid. Biophys J 101:411–420
- Xie J, Schultz PG (2006) A chemical toolkit for proteins—an expanded genetic code. Natl Rev Mol Cell Biol 7:775–782
- Xie J, Supekova L, Schultz PG (2007) A genetically encoded metabolically stable analogue of phosphotyrosine in *Escherichia* coli. ACS Chem Biol 2:474–478
- Young TS, Ahmad I, Yin JA, Schultz PG (2010) An enhanced system for unnatural amino acid mutagenesis in *E. coli.* J Mol Biol 395:361–374

